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(11) EP 0 713 883 B1

(12)

EUROPEAN PATENT SPECIFICATION

- (45) Date of publication and mention of the grant of the patent:
 01.12.1999 Builotin 1999/48
- (21) Application number: 95921149.1
- (22) Date of filing: 09.06.1995

- (51) Int Cl.6: **C08F 10/10**, C08F 2/38, C08F 4/00, C08F 8/00
- (86) International application number: PCT/JP95/01164
- (87) International publication number: WO 95/33774 (14.12.1995 Gazette 1995/53)
- (54) PROCESS FOR PRODUCING ISOBUTENE POLYMER

 VERFAHREN ZUR HERSTELLUNG VON ISOBUTENPOLYMER

 PROCEDE DE PRODUCTION D'UN POLYMERE D'ISOBUTENE
- (84) Designated Contracting States: DE FR GB
- (30) Priority: 09.06.1994 JP 15064694
- (43) Date of publication of application: 29.05.1996 Bulletin 1996/22
- (73) Proprietor: KANEGAFUCHI KAGAKU KOGYO KABUSHIKI KAISHA Kito-ku, Osaka-shi, Osaka 530 (JP)
- (72) Inventors:
 - MAEDA, Takuya
 Tarumi-ku, Kobe-ahi, Hyogo 666 (JP)
 - TAMURA, Masanobu Kobe-shi, Hyogo 654-01 (JP)

- WACHI, Shun
 Takasago-shi, Hyogo 676 (JP)
- (74) Representative: VOSSIUS & PARTNER Siebertstrasse 4 81675 München (DE)
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 DATABASE WPI Section Ch, Week 9046 Derwent Publications Ltd., London, GB; Class A17, AN 90-339137 XP002012265 & JP-A-02 245 004 (KANEGAFUCHI CHEMKK), 28 September 1890

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Description

SPECIFICATION

TECHNICAL FIELD:

[0001] The present invention relates to a process for preparing an isobutene polymer. More particularly, the present invention relates to an advantageous process for preparing an isobutene polymer having functional groups at the terminals in the presence of an initiator.

[0002] The polymer prepared by the process disclosed herein exhibits excellent weathering resistance and gas barder properties and good electrical insulating properties, and thus can be advantageously used as a starting material of coating agent, sealing agent, sealing compound for electronic material, etc.

TECHNICAL BACKGROUND:

[0003] Among telechelic polymers, i.e., polymers having functional groups at a plurality of terminals, polymers having vinyt groups at both terminals are useful as starting materials of photo-setting resins, ultraviolet(UV)-curing reains, electron radiation-curing resins, sealing compounds for electronics, adhesives, modifiers, coating materials, sealing compound for building, etc.

[0004] A functional group-terminated polymer (terminal functional polymers), e.g., isobutone polymor having a chlorine atom bonded to a tertiary carbon at both terminals has been known to be produced by an initiar process which comprises the cationic polymerization of isobutene in the presence of 1,4-bis(α-chloroisopropyl)benzene (hereinafter, simply referred to as "p-DCC") as an initiator/chain transfer agent and boron trichloride as a catalyst (cf. U.S. Patent 4,276,394).

[0005] Further, many roports have been made by Kennedy et al., that when the foregoing cationic polymerization reaction is effected in a solvent containing a halogenated hydrocarbon (e.g., methyl chloride, methylene chloride) in the prosonce of an electron donor, an isobutene polymer having a small Mw/Mn value as determined by GPC, i.e., uniform molecular weight can be obtained (Journal of Macromolecular Science Chemistry, A18 (1), 25 (1982), Polymer Bulletin, 20, 413 (1988), Polymer Bulletin, 26, 305 (1991), JP-A-1-S18014 (the term "JP-A" as used herein means an "unexamined published Japanese patent application")).

[0006] The Inventors made extensive studies of process for the production of an isobutene polymer by the foregoing infer process on a commercial basis. In the course of the studies, the following problems were made clear.

- (1) Since the polymorization reaction is normally effected at low temperatures, the solvent used exhibits a reduced dissolving power. Accordingly, when the monomer concentration is increased to enhance the yield, the resulting polymer separates out. Thus, the monomer concentration can hardly be enhanced. Referring to the process for the proparation of butyl rubber, etc., examples of the preparation process which comprises the procipitation of the resulting polymer have been reported. However, the precipitation of the polymer during the polymerization reaction must be avoided to reduce the molecular weight distribution of the polymer.
 - (2) The polymerization reaction rate is high. Therefore, when the monomer concentration is increased, the resulting heat generation during polymerization makes it difficult to control the reaction temperature. If the rise in the reaction temperature is too great, undesirable side reactions such as proton-inhibited reaction and chain transfer reaction occur, increasing the molecular weight distribution (Mw/Mn) of the polymer and making it difficult to control the terminal functional groups. This disadvantageously produces a mixture of polymers having the following three terminal groups:

Chlorine terminal: -CH₂C(CH₃)₂CI leopropenyl terminal: -CH₂C(CH₃)=CH₂ Internal olefin terminal: -CH=C(CH₃)₂

In particular, if p-DCC is used, the rise in the monomer concentration gives a problem of subsidiary production of a large amount of an Indanyl group having the following structure:

n-C

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(3) Besides the effects on the properties of the polymer thus produced, the problem of toxicity caused by the use of a halogenic solvent arises. Most halogenated hydrocarbons are highly toxic. Halogenated hydrocarbons have recently drawn attention from the standpoint of environmental pollution. Thus, the exhaust of halogenated hydrocarbons out of the system is now under severe regulation. The use of a halogenated hydrocarbon which has heretofore been used in the polymerization reaction, such as methylene chloride and methyl chloride presents problems of deactivation of the catalyst after the polymerization reaction and disposal of waste water or waste solvent containing halogenated hydrocarbon generated at the rinsing step. In order to satisfy the regulation on exhaust, large scale facilities and high disposal cost are required. The regulation will likely be severer in the future, requiring further countermeasures.

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[0007] It has been widely known that when the temperature of the polymerization reaction is raised, carbocations at the growth terminal become instable, causing side reactions such as chain transfer and production-of indanyl group (J.P. Kennedy, E. Marechal, Carbocationic Polymerization, John Wiley & Sons (1992)).

[0008] The rise in the monomer concentration gives a greater rise in the temperature of the polymerization reaction. Therefore, the monomer concentration used in examples which have been reported is low and normally not more than 1 mol/ ℓ (Polym. Bull., 21, 5 (1989), Polym. Bull., 21, 125 (1989), Polym. Bull., 21, 273 (1989), Polym. Bull., 28, 305 (1991), Polym. Bull., 29, 239 (1992)).

[0009] However, from the industrial viewpoint, it has been keenly desired to raise the weight of polymer which can be produced per unit volume.

[0010] Since carbocations as growth seeds must be stable in the polymerization of isobutene in a living cation system, it has heretofore been considered preferred to effect the reaction in a solvent having some polarity at a temperature low enough to allow cations to be present in a relatively stable state. An isobutene polymer exhibits a low solubility in a polar solvent such as methyl chloride and methylene chloride at low temperatures. Thus, a hydrocarbon solvent may be incorporated in the polar solvent to increase the solubility of the isobutene polymer in the polar solvent. However, a report was made that the use of a mixture of methyl chloride and haxane wherein the proportion of haxane la not less than 40% causes chain transfer which appears in the form of two peaks on GPC chart (bimodat) (Polym. Bull., 21, 273 (1989)). The phonomenon that two peaks appear on GPC chart demonstrates the rise in the molecular weight distribution and is thus undestrable.

[0011] Further, JP-A-3-174403, JP-A-3-287605, and U.S. Patente 4,927,201 and 5,189,914 disclose that as a polymerization solvent there may be used an aliphatic organic solvent such as butane, pentane, hexane and heptane, aromatic organic solvent such as benzene, toluene and xylene, nitro compound such as nitromethane and nitrosthane, halogenated organic solvent or mixture thereof without any restriction so far as it exerts no adverse effects on the polymerization activity of the initiator.

[0012] However, the examples in these patents merely give examples of a mixture of methylene chloride and hexane having a high polarity. Further, the resulting isobutene polymer contains a large number of Indanyl groups as by-products. Thus, desirable polymers cannot be obtained. Moreover, as described in the comparative examples of the present application, the polymerization in the presence of hexane, which has a low polarity, produces an isobutene polymer having a molecular weight distribution (Mw/Mn) as extremely high as 7.02 and a large number of indanyl groups. It has thus been made clear that the foregoing polymerization process is not desirable.

40 [0013] An object of the present-invention is to solve the foregoing problems and thus provide a process for the preparation of an industrially favorable isobutene polymer.

DISCLOSURE OF THE INVENTION:

155 [0014] The inventors found that the conventional problems can be solved by the use of a solvent having a specified dielectric constant. The inventors also found that the use of such a solvent makes it possible to obtain the desired isobution polymer having a small molecular weight distribution without using the halogenated hydrocarbon which has heretofore been used. If no halogenated hydrocarbons are used, there arises no problem of the discharge of halogonated hydrocarbons. This advantageously provides a drastic reduction of facilities and cost required for disposal of waste water, making it possible to solve the foregoing problems (1) to (3) at once. However, whether or not such a halogenated hydrocarbon is present has no specific effects on the proporties of the resulting polymer. Even if a solvent containing a halogenated hydrocarbon is used, when the content of halogenated hydrocarbon is reduced to lower the polarity of the solvent, the foregoing problems (1) and (2) can be solved.

[0015] The present invention relates to a process for preparing an isobutene polymor having functional groups at the terminals at a temperature of from not lower than -100°C to not higher than 0°C from the following components as essential components:

(A) a cationically polymerizable monomer containing isobutene;

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(B) an organic compound as an initiator/chain transfer agent represented by formula (1):

$$R^{\frac{1}{2}\left(\begin{array}{c} C - X \end{array} \right)_{n} \tag{1}$$

wherein X represents a halogen atom, an RO- group or an RCOO-group (in which R represents a monovalent organic group); R3 represents a polyvalent aromatic group or a substituted or unsubstituted polyvalent aliphatic hydrocarbon group; R1 and R2 may be the same or different and each represent a hydrogen atom or a substituted or unsubstituted monovalent hydrocarbon group, with the provise that when R3 is a polyvalent aliphatic hydrocarbon group, one of R1 and R2 is not a hydrogen atom; and n represents an integer of from 1 to 6;

(C) an aprotic solvent free of halogenated hydrocarbon; and

(D) a Lewis acid, excluding dialkyl aluminium chloride and dialkyl boron chloride wherein the component (C) exhibits a dielectric constant of from not loss than 2.0 to not more than 4.0 at a temperature of 20°C.

[0016] Furthermore, the invention relates to a process for preparing an isobutane polymer having functional groups at the terminals at a temperature of from not lower than -100°C to not higher than 0°C from the following components as essential components:

(A) a cationically polymerizable monomer containing isobutene;

(B) an organic compound as an initiator/chain transfer agent represented by formula (1):

$$R^{\frac{1}{2}} \begin{pmatrix} R^1 \\ C - X \end{pmatrix}_n \tag{1}$$

wherein X represents a halogen atom, an RO- group or an RCOO-group (in which R represents a monovalent organic group); R³ represents a polyvalent aromatic group or a substituted or unsubstituted polyvalent aliphatic hydrocarbon group; R¹ and R² may be the same or different and each represent a hydrogen atom or a substituted or unsubstituted monovalent hydrocarbon group, with the proviso that when R³ is a polyvalent aliphatic hydrocarbon group, one of R¹ and R² is not a hydrogen atom; and n represents an integer of from 1 to 6;

(C) an aprotic solvent containing helogenated hydrocarbon; and

(D) a Lewis acid, excluding dialkyl aluminium chlorida and dialkyl boron chlorida wherein the component (C) exhibits a dielectric constant of from not less than 3.5 to not more than 5.5 at a temperature of 20°C.

[0017] The component (C) contains preferably an aromatic hydrocarbon.

(0018) The component (C) is preferably a mixture of an aromatic hydrocarbon and an aliphatic hydrocarbon.

[0019] In a proformed embediment the component (C) exhibits a dielectric constant of from not less than 2.0 to not more than 2.5 at a temperature of 20°C.

[0020] The aromatic hydrocarbon can be selected from the group consisting of benzene, toluene, xylene and ethylbenzene.

[0021] The concentration of the component (A) in the system is preferably from 5 to 40 wt%.

[0022] The component (B) may be selected from the group consisting of p-dicumyl chloride, m-dicumyl chloride, p-dicumyl methoxide, m-dicumyl methoxide, 1,3,5-tricumyl chloride and 1,3,5-tricumyl methoxide.

[0023] The component (D) may be boron trichtoride or titanium tetrachlorido.

[0024] An electron donor may further be added as a component (E) in addition to the components (A) to (D).

[0025] The invention relates also to a process for preparing an isobutene polymer having carbon-carbon double bonds at the terminals, which comprises adding an ally! trimethylsiliane in an amount of from 0.5 to 1.5 mol per mol of the group X in the component (B) before the formation of an isobutene polymer terminated by functional groups as described above or before the deactivation of the component (D) following the formation of the isobutene polymer.

[0026] The invention relates also to a process for preparing an isobutene polymer having carbon-carbon double bonds at the terminats, which comprises adding 1,9-decadione in an amount of from 1.5 to 10.0 mol per mol of the

group X in the component (B) before the formation of an isobutene polymer terminated by functional groups as described above or before the deactivation of the component (D) following the formation of the isobutene polymer.

[0027] In the present invention, the number-average molecular weight (Mn) of the isobutene polymer is usually from 500 to 300,000, preferably from 1,000 to 50,000. If Mn is smaller than 500, the excellent characteristics inherent to isobutene polymers are lost. On the contrary, if Mn exceeds 300,000, the resulting polymer is solid, extremely deteriorating its workability. The number-average molecular weight (Mn) and Mw/Mn value of the isobutene polymer were determined by GPC using a polystyrene gel column (Shodex K-804, manufactured by Showa Denko K.K.; mobile phase: chloroform) in polystyrene equivalence.

[0028] The term "cationically polymerizable monomer containing isobutene" as used herein is not limited to a monomer made only of isobutene but is meant to include a monomer the isobutene component of which is substituted by a cationically polymerizable monomer copolymerizable with isobutene in a proportion of not more than 50% by weight (hereinafter, simply referred to as "%").

[0029] Examples of the cationically polymerizable monomer copolymerizable with isobutene include olefins having 3 to 12 carbon atoms, conjugated dienes, vinyl ethers, aromatic vinyl compounds, norbomenes, and vinyl silanes. Proferred among these cationically polymerizable monomers are olefins having 3 to 12 carbon atoms and aromatic vinyl compounds.

[0030] Specific examples of the cationically polymerizable monomer copolymerizable with isobutene usually include propene, 1-butene, 2-methyl-1-butene, 3-methyl-1-butene, pentene, hexene, cyclohexene, vinyl cyclohexene, 5-ethylidonenorbornene, 5-propylidonenorbornene, butadiene, isoprene, cyclopentadiene, methyl vinyl ether, ethyl vinyl ether, leobutyl vinyl ether, vinyl carbazole, methoxystyrene, ethoxystyrene, t-butoxystyrene, hexenyloxystyrene, styrene, α-methylstyrene, methylstyrene, dimethylstyrene, chloromethylstyrene, chlorostyrene, indene, β-pinene, vinyltrichlorosilane, vinylmethyldichlorosilane, vinyldimethylchlorosilane, vinyldimothylmothoxysilane, vinyltrimethylsilane, divinyldichlorosilane, divinyldimethoxysilane, divinyldimethylsilane, γ-methacryloyloxypropyltrimethoxysilane, and γ-methacryloyloxypropylmothyldimethoxysilane.

[0031] Preferred among these cationically polymerizable monomers are propene, 1-butene, 2-butene, cyclopentadiene, 5-ethylidenenorbomene, isobutyl vinyl ether, methoxystyrene, and styrene. One of these cationically polymerizable monomers copolymerizable with isobutene may be used in combination with isobutene. Two or more of these cationically polymerizable monomers may be used in combination.

[0032] In the present invention, the monomer concentration during polymerization is normally from 1 to 75 wt%, proferably from 5 to 60 wt%, more preferably from 10 to 40 wt%, most proferably from 15 to 30 wt%. If the monomer concentration is less than 1 wt%, the production efficiency is disadvantageously reduced, though causing no troubles on the polymerization litself. If the monomer concentration exceeds 75 wt%, the polymerization system is a precipitation system, making it impossible to obtain a good polymer.

[0033] Examples of the compound represented by the foregoing formula (1) of the present invention include a compound represented by formula (2):

40 wherein A represents a group having 1 to 4 aromatic rings; Y represente a group represented by formula (3):

$$\begin{array}{c}
R^6 \\
 \downarrow \\
-C-X \\
\downarrow \\
R^7
\end{array}$$
(3)

(in which R⁶ and R⁷ may be the same or different and each represent a hydrogen atom or a monovalent hydrocarbon group having 1 to 20 carbon atoms; X represents a halogen atom, an R⁴COO- group (R⁴ represents a hydrogen atom or an alkyl group having 1 to 5 carbon atoms) or an R⁶O- group (R⁵ represents a hydrogen atom or an alkyl group having 1 to 5 carbon atoms)); and n represents an integer of from 1 to 8; a compound represented by formula (4):

wherein B represents a substituted or unsubstituted hydrocarbon group having 4 to 40 carbon atoms; Z represents a halogen atom bonded to tertiary carbon atom, an P⁸COO- group (in which R⁸ represents a hydrogen atom or an alkyl group having 1 to 5 carbon atoms) or an R⁹O- group (in which R⁹ represents a hydrogen atom or an alkyl group having 1 to 5 carbon atoms); and m represents an integer of from 1 to 4; and an oligomer having α-halostyrene units. However, the present invention is not limited to these compounds. These compounds may be used singly or in combination.

[0034] In the compound represented by the formula (2), the group A having 1 to 4 aromatic rings may be either one produced by a condensation reaction or may be an uncondensed product. Examples of the group containing aromatic rings include phenyl, biphenyl, naphthyl, anthryl, phenanthryl, pyrenyl, di- to pentavalent groups derived from these groups, and mono- to hexavalent groups derived from Ph-(CH₂)₁-Ph (in which Ph represents a phenyl group, and 1 represents an integer of from 1 to 10). These groups having aromatic rings may be substituted by straight-chain and/ or branched allphatic hydrocarbon groups having 1 to 20 carbon atoms or groups containing a functional group such as hydroxyl group, other group and vinyl group.

[0035] As the compound represented by the formula (4), there may also be used one having a functional group other than Z, such as vinyl group and sityl group.

[0036] Examples of the oligomer having an α -haloetyrene unit, which can be used as the initiator/chain transfer agent, include oligomers of α -chloroetyrene, and oligomers obtained by the copolymerization of α -chloroetyrene with a monomer copolymerizable therewith.

[0037] In the present invention, the use of a compound having two or more halogen atoms, R4COO- groups (in which R4 represents a hydrogen atom or an alkyl group having 1 to 5 carbon atoms) or R5O- groups (in which R5 represents a hydrogen atom or an alkyl group having 1 to 5 carbon atoms) or a compound having a halogen atom, an R4COO-group or an R5O- group and other reactive functional groups, from among the compounds represented by the formula (1), as an initiator/chain transfer agent, can advantageously provide the resulting polymer with a higher functionality. [0038] Specific examples of the compound represented by the foregoing formula (1), which can be usually employed, include the following compounds:

CX(CH₂)₂

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and

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wherein X represents a halogen atom, an R⁴COO- group (in which R⁴ represents a hydrogen atom or an alkyl group having 1 to 5 carbon atoms) or an R⁵O- group (in which R⁵ represents a hydrogen atom or an alkyl group having 1 to 5 carbon atoms); and

oligomer of α -chlorostyrene. However, the present invention is not limited to these compounds. Preferred among these compounds are compounds such as

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and CH₃O- group-containing compounds such as

 $CH_{3} CH_{3} CH_{3}$ $CH_{3} CH_{3} CH_{3} CH_{3} CH_{3} CH_{3} CH_{3} CH_{3} CH_{3}$

and

 $cH_{3}O - c C(cH_{3})_{2}OcH_{3}$

[0039] Those compounds are components which are used as initiators/chain transfer agents. These compounds may be used singly or in admixture in the present invention. By controlling the amount of these compounds to be used, the number-average molecular weight of the resulting isobutone polymer can be arbitrarily predetermined.

[0040] In the present invention, the amount of the compound represented by the foregoing formula (1) to be used is usually from 0.01 to 20% by weight, preferably from 0.1 to 10% by weight based on the weight of the cationically polymerizable monomer containing isobutene.

[0041] As the Lewis acid which can be used in the present invention, there may be used any known metal complex apart from dialkyl aluminium chloride and dialkyl boron chloride. In general, TiCl₄, BCl₃, SnCl₄, AlCl₃, CH₃CH₂AlCl₂, VCl₅, FeCl₃, BF₃, etc. may be used. However, the present invention is not limited to those compounds. Preferred among these Lewis acids are TiCl₄ and BCl₃.

[0042] In the present invention, the Lewis acid is usually used in an amount of from 0.1 to 100% by weight, preferably from 1 to 30% by weight based on the weight of the cationically polymerizable monomer containing isobutene. If the amount of the Lewis acid to be used as a catalyst component falls below 0.1% by weight, a problem arises that the yield of the product of the cationic polymerization reaction is reduced. On the contrary, if the amount of the Lewis acid exceeds 100% by weight, no morits can be obtained.

[0043] In the present invention, the aprotic solvent free of halogenated hydrocarbon, if used as a polymerization

solvent, should have a dielectric constant of from not less than 2.0 to not more than 4.0 at a temperature of 20°C. If the dielectric constant of the solvent falls below 2, a good isobutene polymer could not be obtained. (Therefore, pentane, hexane, etc. cannot be used as single solvent.)

[0044] Examples of the aprotic solvent free of halogenated hydrocarbon to be used as a polymerization solvent include butane, pentane, neopentane, hexane, cyclohexane, methylcyclohexane, ethylcyclohexane, heptane, octane, benzene, toluene, xylene, and ethylbenzene. Professed among these aprotic solvents are toluene, hexane, and heptane.

[0045] Examples of the mixed solvent (mixing ratio (volvol)) include various combinations such as toluene/hexane (5/5 to 10/0), toluene/heptane (5/5 to 10/0), toluene/heptane (5/5 to 10/0), toluene/heptane (5/5 to 10/0). One of unexpected merits which have been made obvious in the course of the study in the present invention is the fact that the use of a solvent free of halogenated hydrocarbon makes it possible to obtain a good isobutene polymer even if the dielectric constant of the solvent is low. A low dielectric constant system tends to show a small temperature rise during polymerization reaction. Therefore, the problem of heat removal accompanied by scaling up the reaction can be easily solved.

[0046] In the present invention, the aprotic solvent containing halogenated hydrocarbon, if used as a polymerization solvent, should have a dielectric constant of from not less than 3.5 to not more than 5.5 at a temperature of 20°C.

[0047] Specific examples of the aprotic solvent containing halogenated hydrocarbon to be used as a polymerization solvent include a mixture of halogenated hydrocarbon such as methyl chloride, methylene chloride, 1,2-dichloroethane, 1,1-dichloroethane and chlorination product of propane, butane and pentane and hydrocarbon such as butane, pentane, neopentane, hexane, cyclohexane, methylcyclohexane, ethylcyclohexane, heptane, octane, benzene, folluene and xytene. Examples of the mixed solvent include various combinations such as methylene chloride/hexane, methyl chloride/hexane, and methylene chloride/heptane. Heretofore, if a solvent containing halogenated hydrocarbon is used, the polymerization has been often effected in a solvent system having a dielectric constant of not less than 6. In the present invention, by reducing the dioloctric constant of the solvent, the polymerization rate can be controlled, making

it possible to control the temporature rise during polymerization.

[0048] In the present invention, the dielectric constant of the mixed solvent system can be approximately calculated by determining the additive mean (arithmetic mean) of the dielectric constant of the constituents over the mixing ratio of the constituents on a weight-average basis. The dielectric constant (typical value) of various solvents used for the calculation of the dielectric constant of the mixed solvent will be listed below.

30 [0049] Tolueno: 2.24; ethylbenzene: 2.40; o-xylene: 2.27; m-xylene: 2.37; n-pentane: 1.84; n-hexane: 1.89; n-heptane: 1.92; methylcyclohexane: 2.02; methylene chlorido: 7.77; methyl chloride: 12.9

[0050] In the present invention, the polymerization solvent is preferably free of halogenated hydrocarbon for the reason that waste water disposal facilities can be simplified. The term "solvent free of halogenated hydrocarbon" as used in the present application is meant to indicate a solvent which may contain halogenated hydrocarbon to an extent such that no special treatments for halogenated hydrocarbon in waste water or waste solvent generated from the catalyst deactivation and dhalog process after polymerization reaction are required.

[0051] In the present invention, an electron donor is preferably used as the component (E) to control side reactions more efficiently. As the electron donor component there may be widely used any known electron donor components having a donor number of from 15 to 50. Preferred examples of such electron donor components include pyridines, amines, and sulfoxides.

[0052] Specific examples of the electron donor component usually include 2,6-dirt-butylpyridine, 2-t-butylpyridine, 2,4,6-trimethylpyridine, 2,6-dimethylpyridine, 2-methylpyridine, pyridine, diethylamine, trimethylamine, triethylamine, tributylamine, diethylamine, diethylamine, N,N-dimethylamine, N,N-dimethylamine, trimethylamine, N,N-dimethylamine, N,N-dimethylamine, N,N-dimethylamine, N,N-dimethylamine, and triamide hoxamethylphosphate. Preferred among these electron donor components are 2,6-di-t-butylpyridine, 2,6-dimethylpyridine, 2-methylpyridine, pyridine, diothylamine, trimethylamine, triethylamine, N,N-dimethylformamide, N,N-dimethylamide, and dimethylsulfoxide. Picolines are further preferred. Particularly preferred of these electron donors is a-picoline (2-moth-

ylpyridine), which can exert a remarkable effect despite of its small donor number.

[0053] In the present invention, the electron donor component is usually used in an amount of from 0.01 to 10 mol, preferably from 0.1 to 2 mol per mol of the group X in the compound represented by the general formula (1).

[0054] In the present Invention, the polymerization reaction is not specifically limited and can be accomplished by any conventional polymerization method. Among the foregoing components, i.e., (A) cationically polymerizable monomer containing isobutene and (D) Lewis acid, a solution containing (D) a Lewis acid may be added to a solution of (A) a cationically polymerizable monomer containing isobutene in a batch process. Alternatively, a solution of (A) a cationically polymerizable monomer containing isobutylene may be continuously added to a solution containing (D) a Lewis acid in a semi-batch process. Further, a continuous process may be employed which comprises continuously charging all the foregoing components into the reaction system while withdrawing the reaction product.

[0055] In the present invention, the polymorization reaction can be followed by a substitution reaction or addition

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reaction on the functional terminal groups formed at the terminals of the polymer to produce an isobutene polymer having carbon-carbon double bonds at the terminals. In other words, the reaction of the resulting isobutene polymer, preferably at tertiary chlorine group end, with a specific end capping agent makes it easy to obtain a polymer having carbon-carbon double bonds at the terminals. In general, the functional terminal group is chiefly a tertiary halogen group. However, the foregoing isopropenyl group or internal olefin group may be included.

[0056] For example, in the conversion of chlorine group end to carbon-carbon double bond end by the substitution reaction using an allyl trimethylsiliane, the desired carbon-carbon double bond end can be easily produced only by adding a required amount of an allyl trimethylsiliane before the deactivation of the component (D). The reaction temperature is preferably as low as the polymerization temperature. Since the substitution reaction proceeds at a high rate at low temperatures, it is usually completed in 30 minutes. The amount of the allyl trimethylsiliane to be used may be from 1.0 to 1.5 mol per mol of the group X in the component (B). If it is necessary to obtain a polymer having a small content of carbon-carbon double bonds, the amount of the allyl trimethylsiliane added may be not more than 1 mol per mol of the group X. Since the allyl trimethylsiliane doesn't take part in the polymorization reaction, it may be added to the reaction system before polymerization.

[Reaction scheme]

[0057]

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P-C(CH₃)₂CI → P-C(CH₃)₂CH₂CH=CH₂

wherein P represents a polymer residue.

[0058] Further, in the conversion of chlorino group and to carbon-carbon double bond end by the addition reaction using a non-conjugated diene such as 1,9-decadiene, the desired carbon-carbon double bond end can be easily produced only by adding a required amount of a non-conjugated diene before the deactivation of the component (D) as in the case where an allyl trimethylsilane is used. The addition reaction temperature is preferably as low as the polymerization temperature. The addition reaction at low temperatures is usually completed in about 6 hours. The amount of the non-conjugated diene to be used may be from 1.5 to 10.0 mol per mol of the group X in the component (B). If it is necessary to obtain a polymer having a small carbon-carbon double bond content, the reaction time may be reduced. Alternatively, the amount of 1,9-decadiene added may be not more than 1 mol per mol of the group X. In this system, a Lewis acid as the component (D) may be added, or the reaction temperature may be further lowered for the purpose of further reducing the addition reaction rate.

[Reaction scheme]

[0059]

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P-C(CH₃)₂Cl → P-C(CH₃)₂CH₂CHCl(CH₂)₆CH=CH₂

wherein P represents a polymer residue.

BEST MODE FOR CARRYING OUT THE INVENTION:

[0060] The present invention will be further described in the following examples, but the present invention should not be construed as being limited thereto.

Example 1:

[0061] In a reaction vessel were charged 276 mℓ of toluene, 112 mℓ of an isobutene monomer, 1.156 g of p-DCC and 0.186 g of α-picoline. The reaction mixture was then stirred with the reaction vessel being surrounded by a dry ice-ethanol bath so that it was cooled to a temperature of -70°C. A mixture of 3.95 mℓ of TiCl₄ and 4 mℓ of toluene was then added to the reaction system to initiate reaction. After the completion of the reaction, the reaction solution was poured into a large amount of water with stirring so that it was washed. The resulting organic phase and aqueous phase were then separated from each other to remove the catalyst. The organic phase was then subjected to evaporation under reduced pressure to remove the volatile content. Thus, an isobutene polymer was obtained. The reaction conditions used and the properties of the polymer thus obtained are shown in Tables 1 and 2 below.

Example 2:

[0062] The air in a 500-m² four-necked flack with a throo-way cock was replaced by nitrogen. Into the vossel were charged 280 m² of toluene (which had been allowed to stand with molecular sieves 3A overnight or longer so that it was dried) and 2.91 g (10 mmol) of p-DCC through a syringe. To the three-way cock was then connected a pressure glass liquefied gas thief tube equipped with a needle valve in which 112 m² of an isobutene monomer had been charged. The polymerization vessel was then dipped in a -60°C dry los/ethanol bath so that it was cooled. The system in the vassel was then evacuated by a vacuum pump. The needle valve was then opened so that the isobutene monomor was introduced from the liquefied gas thiof tube into the polymerization vessel. Nitrogen gas was then introduced into the polymorization vessel from one of the three ways of the cock so that the pressure of the vessel was returned to normal value (atmospheric pressure). 0.186 g (2 mmol) of 2-methylpyridine was then charged into the polymerization vessel. 2.74 m² (25 mmol) of titanium tetrachloride was then added to the -60°C monomer solution to initiate polymerization. When 60 minutes passed since the initiation of the reaction, 3.42 g (30 mmol) of an allyl trimethylsilane was added to the reaction mixture. The reaction mixture was further allowed to undergo reaction at a temperature of -60°C for 120 minutes. The reaction solution was then washed with 200 m² of water four times. The solvent was then evaporated to obtain an isobutene polymer. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2.

Example 3:

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[0083] The air in a 300-m/ four-necked flask with a three-way cock was replaced by nitrogen. Into the vessel were charged 112 m/ of toluene (which had been allowed to stand with molecular sieves SA overnight or longer so that it was dried), 49 m/ of heptane (which had been allowed to stand with molecular sieves SA overnight or longer so that it was dried), 1.16 g (5.0 mmol) of p-DCC and 0.093 g (1 mmol) of 2-methylpyridino through a syringe. To the three-way cock was then connected a pressure glass liquefled gas thlef tube equipped with a needle valve in which 56 m/ of an isobutene monomer had been charged. The polymorization vessel was then dipped in a -70°C dry ico/othanol bath so that it was cooled. The system in the vessel was then evacuated by a vacuum pump. The needle valve was then opened so that the isobutene monomer was introduced from the liquefled gas thief tube into the polymerization vessel. Nitrogen gas was then introduced into the polymerization vessel from one of the three ways of the cock so that the pressure of the vessel was returned to normal value. 1.64 m/ (15 mmol) of titanium tetrachloride was then added to the monomer solution to initiate polymerization. When 90 minutes passed since the initiation of the reaction, 1.15 g (10 mmol) of an allyl trimethylsilane was added to the reaction mixture. The reaction mixture was further allowed to undergo reaction at a temperature of -70°C for 90 minutes. The reaction solution was then washed with 200 m/ of water four times. The solvent was then evaporated to obtain an isobutene polymer. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2.

Example 4:

[0064] An isobutene polymer was produced in the same manner as in Example 3 except that 128 mf of follower and 32 mf of heptane were used as solvents. The isobutene polymer thus produced was then evaluated in the same manner as in Example 3. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2. Example 5:

[0065] The air in a 500-m² four-necked flask with a three-way cock was replaced by nitrogen. Into the vessel were charged 255 mf of toluene (which had been allowed to stand with molecular eleves 3A overnight or langer so that it was dried), 112 mf of hexane (which had been allowed to stand with molecular sleves 3A overnight or longer so that it was dried) and 1.16 g (5.0 mmol) of p-DCC through a syringe. To the three-way cock was then connected a pressure glass liquefied gas thief tube equipped with a needle valve in which 112 m² of an isobutene monomer had been charged. The polymerization vessel was then dipped in a .70°C dry ice/othanol bath so that it was cooled. The system in the vessel was then evacuated by a vacuum pump. The needle valve was then opened so that the isobutene monomer was introduced from the liquefied gas thief tube into the polymerization vassal. Nitrogen gas was then introduced into the polymerization vessel from one of the three ways of the cock so that the pressure of the vessel was returned to normal value, 0,188 g (2 mmol) of 2-methylpyridine was then added to the reaction system. 7 mf of a toluene solution of 2.76 mg (25 mmol) of titanium tetrachloride was then added to the reaction system to initiate polymerization. When the reaction was initiated, the temperature rise was 6.5°C. When 2 hours passed since the initiation of the reaction, 5.48 mf (50 mmol) of titanium tetrachloride and 13.83 g (100 mmol) of 1,9-decadiene were added to the reaction system. When 6 hours passed since the addition of 1,9-decadens, the reaction solution was washed with 500 mf of water four times. The solvent was evaporated. The resulting isobutone polymer was then dissolved in a small amount of hexane. 1,000 mf of acetone was then added to the reaction solution with stirring so that the polymer separated out

by the precipitation. The polymer thus obtained was dissolved in hexano. The volatile content was then evaporated to obtain an isobutone polymer. The reaction conditions used and the properties of the resulting polymer are attogether shown in Tables 1 and 2.

Example 6:

[0066] The air in a 500-mf four-necked flask with a three-way cock was replaced by nitrogen. Into the vessel were charged 126 mf of toluene (which had been allowed to stand with molecular sieves 3A overnight or longer so that it was dried), 54 mf of methyloyclohoxane (which had been allowed to stand with molecular sieves 3A overnight or longer so that it was dried) and 1.16 g (5.0 mmol) of p-DCC through a syringe. To the throe-way cock was then connected a pressure glass liquofied gas thiof tube equipped with a needle valve in which 56 mf of an isobutene monomer had been charged. The polymerization vessel was then dipped in a -70°C dry ice/ethanol bath so that it was cooled. The system in the vessel was then evacuated by a vacuum pump. The needle valve was then opened so that the isobutone monomer was introduced from the liquefied gas thief tube into the polymerization vessel. Nitrogen gas was then introduced into the polymerization vessel from one of the three ways of the cock so that the pressure of the vessel was returned to normal value. 0.093 g (1 mmol) of 2-methylpyridine was then added to the reaction system. 1.65 mf (15.1 mmol) of titanium totrachloride was then added to the reaction system.

[0067] When 70 minutes passed since the initiation of the reaction, 1.24 g (10.8 mmol) of an allyl trimethylsitane was added to the reaction system. The reaction system was further allowed to undergo reaction at a temperature of -70°C for 90 minutes. The reaction solution was then washed with 400 mf of water four times. The solvent was evaporated. The resulting isobutene polymer was then dissolved in a small amount of hexane. 500 mf of acetone was then added to the solution with stirring so that the polymer separated out by the precipitation. The polymer thus obtained was dissolved in hexane. The volatile content was then evaporated to obtain an isobutene polymer. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2.

Example 7:

[0068] An Isobutene polymer was produced in the same manner as in Example 1 except that 28 mf of an Isobutene monomor, 0.289 g of ρ-DCC and 0.047 g of α-picoline were charged into the reaction system. The Isobutene polymer thus produced was then evaluated in the same manner as in Example 1. The reaction conditions used and the properties of the polymer thus obtained are altogether shown in Tables 1 and 2. The maximum temperature rise during reaction was 5°C.

Example 8:

[0069] An isobutene polymer was produced in the same manner as in Example 7 except that the polymerization temperature was -50°C. The isobutone polymer thus produced was then evaluated in the same manner as in Example 7. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2.

40 Example 9:

[0070] An isobutene polymer was produced in the same manner as in Example 7 except that 248 m/ of toluene and 28 m/ of hexane were charged into the reaction vessol. The isobutene polymer thus produced was then evaluated in the same manner as in Example 7. The reaction conditions used and the properties of the polymer thus obtained are altogether shown in Tables 1 and 2.

Comparative Example 1:

[0071] Into a reaction vessel were charged 276 mℓ of hexane, 28 mℓ of an isobutene monomer, 0.289 g of ρ-DCC and 0.047 g of α-picoline. The reaction mixture was then altired with the reaction vessel being surrounded by a dry ice-ethanol bath so that it was cooled to a temperature of -70°C. A mixture of 7.9 mℓ of TiCl₄ and 4 mℓ of hexane was then added to the reaction system to initiate reaction. After the completion of the reaction, the reaction solution was poured into a large amount of water with stirring so that it was washed. The resulting organic phase and aqueous phase were then separated from each other to remove the catalyst. The organic phase was then subjected to evaporation under reduced pressure to remove the volatile content. Thus, an isobutene polymer was obtained. The reaction conditions used and the properties of the polymer thus obtained are altogether shown in Tables 1 and 2 below.

Example 10:

[0072] Into a reaction vessel were charged 76 mf of methane dichloride, 196 mf of hoxane, 28 mf of an isobutene monomer, 0.289 g of p-DCC and 0.047 g of α-picoline. The reaction mixture was then stirred with the reaction vessel being surrounded by a dry los-ethanol bath so that it was cooled to a temperature of -70°C. A mixture of 3.95 mf of TiCl₄ and 8 mf of methane dichloride was then added to the reaction system to initiate reaction. After the completion of the reaction, the reaction solution was poured into a large amount of water with stirring so that it was washed. The resulting organic phase and aqueous phase were then separated from each other to remove the catalyst. The organic phase was then subjected to evaporation under reduced pressure to remove the volatile content. Thus, a polymer product was obtained. The reaction conditions used and the properties of the polymer thus obtained are altogether shown in Tables 1 and 2 below.

Example 11:

[0073] Into a reaction vessel were charged 56 mf of methane dichloride and 216 mf of hexane. The other reaction conditions were the same as used in Example 10. A solution of 3.95 mf of a catalyst in 8 mf of hexane was then poured into the reaction vessel to initiate reaction. Thus, a polymer product was obtained. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2 below. The maximum temperature rise during reaction was 7°C.

Example 12:

[0074] A reaction was effected in the same manner as in Example 10 except that 104 mt of methane dichloride and 168 mt of hexane were charged into the reaction vessel. Thus, a polymer product was obtained. During the reaction, 9.95 g of 1,9-decadiene was added to the reaction system to introduce a functional group into the polymer. The reaction temperature was -50°C. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2 bolow.

[0075] The molecular weight of the polymer product and its distribution were measured by GPC analysis. As a result, no low molecular by-products were detected. The maximum temperature rise during reaction was 7°C.

Comparative Example 2:

[0076] Into a reaction vessel were charged 28 mf of methane dichloride and 244 mf of hoxane. The other conditions were the same as used in Example 10. A solution of 3.95 mf of a catalyst in 8 mf of hexane was then poured into the reaction vessel to initiate reaction. Thus, a polymer product was obtained. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2 below.

Comparative Example 3:

[0077] A reaction was effected in the same manner as in Example 12 except that 160 mf of methane dichloride, 112 mf of hexane and 0.578 g of p-DCC were charged into the reaction vossel. Thus, a polymer product was obtained. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2 below. The molecular weight of the polymer product and its distribution were measured by GPC analysis. As a result, low molecular by-products were detected in a molar fraction of 0.2. The maximum temperature rise during reaction was 9°C.

Comparative Example 4:

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[0078] A reaction was effected in the same manner as in Comparative Example 3 except that 188 mt of methane dichloride and 84 mt of hexane were charged into the reaction vessel. Thus, a polymer product was obtained. The reaction conditions used and the properties of the resulting polymer are altogether shown in Tables 1 and 2 below. The molecular weight of the polymer product and its distribution were measured by GPC analysis. As a result, low molecular by-products were detected in a molar fraction of 0.4. The maximum temperature dee during reaction was 10°C.

Table 1

•		Solvent (charged amount (mf))	Diolectric Constant	Monomer Concentration (wl%)
	Example 1	toluono (276)	2.2	24 .

Table 1 (continued)

		Solvent (charged amount (m/))	Dielectric Constant	Monomer Concentration (wt%)	
5	Example 2	toluene (280)	2.2	24	
	Examplo 3	toluene (112), heplane (49)	2.2	36	
10	Example 4	toluono (128), heptane (32)	2.2	36	
	Example 5	toluene (262), hexane (112)	2.2	20	
15	Example 6	toluena (126), methylcyclohexane (54)	2.2	20	
	Example 7	toluene (276)	2.2	7	
20	Example 8	Example 8 toluena (276)		7	
	Example 9	toluene (248), haxane (28)	2.2	ə	
25	Example 10	mothylene chloride (84), hexane (196)	4.6	. 4	
	Examplo 11	mothylene chloride (56), 3.9 hexane (224)		a .	
30	Example 12	melhylene chloride (112), hoxane (168)	5,0	7	
35	Comparative Example 1	hexane (276)	1.9	· 9	
	Comparative Example 2	methylene chloride (28), hexane (252)	3,0	В	
40	Comparative Example 3	methylene chlorido (168), hoxane (112)	6.3	6	
	Comparativa Example 4	melhylene chloride (196), hexane (84)	6.7	6	

[0079] It can be seen from the results shown in Table 1 that Examples 1 to 9, wherein an aprotic solvent free of halogenated hydrocarbon was used as a polymerization solvent, exhibit a dielectric constant of from not less than 2.0 to not more than 4.0. It can also be seen that Examples 10 to 13, wherein a solvent containing halogenated hydrocarbon was used, exhibit a dielectric conclant of from not less than 3.5 to not more than 5.5.

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Table 2

	GPC		NMR		
	Mn	Mw/Mn	Fn(C=C)1)	Fn(indanyl)2)	Mn(cal.)
Example 1	16,000	1.13	•	0.16	
Example 2	7,200	1.24	1.95	0.0	9,100
Example 9	7,700	1.38	2.09	0.0	8,700
Example 4	7,900	1.32	2.02	0.0	8,800
Example 5	16,700	1.27	1.84	0.0	19,500
Example 6	8,800	1.37	1.90	0.0	9,600
Example 7	14,600	1.12	•	0.17	-
Example 8	14,300	1.2	•	0.1B	-
Example 9	16,100	1.10	•	0.14	
Example 10	18,500	1.07	-	0.11	
Example 11	15,800	1.08	-	0.13	
Example 12	18,500	1.2	1.79	0.17	•
Comparativo Example 1	1,550	7.02	•	0.87	1 •
Comparativo Example 2	2,060	7.44	-	03,0	
Comparative Example 3	9,310	1.75	•	0.21	•
Comparative Example 4	9,270	1.6	•	0.26	•

Note

2) Average by number of indanyl groups contained in one molecule of isobutone polymer

[0080] The results in Table 2 show that a solvent system having a high monomer concentration free of halogenated hydrocarbon can provide an isobutene polymer having a low Mw/Mn ratio and a high terminal functionality. It can be also seen from the results of Comparative Example 1 that a solvent system comprising haxane alone cannot provide a good polymer.

INDUSTRIAL APPLICABILITY:

[0081]

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- (1) By selecting a specific solvent system, it is made possible to prevent the resulting polymer from being precipitated and enhance the monomer concentration during polymerization.
- (2) By selecting a specific solvent system, it is made possible to obtain a good polymor having a narrow molecular weight distribution and a small number of indanyl groups.
- (3) If a solvent free of halogenated hydrocarbon is used, no special treatments for halogenated hydrocarbon in waste water or waste solvent generated from the catalyst deactivation and rinsing process after polymerization reaction are required.

45 Claims

- A process for preparing an isobutene polymer having functional groups at the terminals at a temperature of from not lower than -100°C to not higher than 0°C from the following components as essential components:
 - (A) a cationically polymerizable monomer containing isobutene;
 - (B) an organic compound as an initiator/chain transfer agent represented by formula (1):

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¹⁾ Average by number of carbon-carbon double bonds continined in one molecule of leobutane polymer

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$$R^{2} = \left(\begin{array}{c} R^{1} \\ C \\ -X \end{array} \right)_{n} \tag{1}$$

wherein X represents a hatogen atom, RO- group or RCOO- group (in which R represents a monovalent organic group); R⁹ represents a polyvalent aromatic group or substituted or unsubstituted polyvalent aliphatic hydrocarbon group; R¹ and R² may be the same or different and each represent a hydrogen atom or substituted or unsubstituted monovalent hydrocarbon group, with the proviso that when R⁹ is a polyvalent aliphatic hydrocarbon group, one of R¹ and R² is not a hydrogen atom; and n represents an integer of from 1 to 6;

- (C) an aprotic solvent free of halogonated hydrocarbon; and
- (D) a Lewis acid, excluding dialkyl aluminium chloride and dialkyl boron chloride.

wherein said component (C) exhibits a dielectric constant of from not less than 2.0 to not more than 4.0 at a temperature of 20°C.

- A process for preparing an isobuteno polymer having functional groups at the terminals at a temporature of from not lower than -100°C to not higher than 0°C from the following components as essential components:
 - (A) a cationically polymerizable monomer containing isobutene;
 - (B) an organic compound as an initiator/chain transfer agent represented by formula (1):

$$R^{\frac{1}{2}} \left(\begin{array}{c} R^{1} \\ C - \times \end{array} \right)_{R} \tag{1}$$

wherein X represents a halogen atom, RO- group or RCOO- group (in which R represents a monovalent organic group); R3 represents a polyvalent aromatic group or substituted or unsubstituted polyvalent aliphatic hydrocarbon group; R1 and R2 may be the same or different and each represent a hydrogen atom or substituted or unsubstituted monovalent hydrocarbon group, with the proviso that when R3 is a polyvalent aliphatic hydrocarbon group, one of R3 and R2 is not a hydrogen atom; and n represents an integer of from 1 to 6;

- (C) an aprotic solvent containing halogenated hydrocarbon; and
- (D) a Lewis acid, excluding dialkyl aluminium chloride and dialkyl boron chloride
- wherein said component (C) exhibits a dielectric constant of from not less than 3.5 to not more than 5.5 at a temperature of 20°C.
 - The process for preparing an isobutene polymer terminated by functional groups according to Claim 1 or 2, wherein said component (C) contains an aromatic hydrocarbon.
 - 4. The process for preparing an isobutene polymer terminated by functional groups according to any one of Claims 1 to 3, wherein said component (C) is a mixture of an aromatic hydrocarbon and an aliphatic hydrocarbon.
- 5. The process for preparing an isobutone polymer terminated by functional groups according to any one of Claims
 1, 3 and 4, wherein said component (C) exhibits a dielectric constant of from not less than 2.0 to not more than
 2.5 at a temperature of 20°C.
 - The process for preparing an isobutene polymer terminated by functional groups according to any one of Claims 3 to 5, wherein said aromatic hydrocarbon is selected from the group consisting of benzene, toluene, xylone and ethylbenzene.
 - The process for preparing an isobutene polymer terminated by functional groups according to any one of Claims 1 to 6, wherein the concentration of said component (A) in the system is from 5 to 40 wt%.

- 8. The process for preparing an isobutene polymer terminated by functional groups according to any one of Claims 1 to 7, wherein said component (B) is selected from the group consisting of p-dicumyl chloride, m-dicumyl methoxide, 1,3,5-tricumyl chloride and 1,3,5-tricumyl methoxide.
- 5 9. The process for preparing an isobutone polymer terminated by functional groups according to any one of Claims
 1 to 8, wherein said component (D) is boron trichloride or titanium tetrachloride.
 - 10. The process for preparing an isobutene polymer terminated by functional groups according to any one of Claims 1 to 9, wherein an electron donor is further added as a component (E) in addition to said components (A) to (D).
 - 11. A process for preparing an isobutene polymor having carbon-carbon double bonds at the terminals, which comprises adding an allyl trimethylsilane in an amount of from 0.5 to 1.5 mol per mol of the group X in said component (B) before the formation of an isobutene polymer terminated by functional groups according to any one of Claims 1 to 10 or before the deactivation of said component (D) following the formation of said isobutene polymer.
 - 12. A process for preparing an isobutene polymer having carbon-carbon double bonds at the terminals, which comprises adding 1,9-decadiene in an amount of from 1.5 to 10.0 mol per mol of the group X in said component (B) before the formation of an isobutene polymer terminated by functional groups according to any one of Claims 1 to 10 or before the deactivation of said component (D) following the formation of said isobutene polymer.

Patentansprüche

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- 1. Verfahren zur Hersteilung eines Isobutenpolymers, das endständige funktionelle Gruppen aufwelst, bei einer Temperatur nicht unterhalb von -100°C bis nicht oberhalb von 0°C aus den folgenden Komponenten als wesentlichen Komponenten:
 - (A) ein kationisch polymerisierbares Monomer, das Isobuten enthält;
 - (B) eine organische Verbindung als Initiator/Kettenübertragungsreagens, dargestellt durch die Formel (I)

$$8^{2} + \left(\begin{array}{c} 2 \\ 0 \\ 0 \end{array} \right)_{\pi}$$

wobei X ein Halogenatom, einen RO-Rest oder RCOO-Rest darstellt (wobei R einen einwertigen organischen Rest darstellt); R³ einen mehrwartigen aromatischen Rest oder einen substituierten oder unsubstituierten mehrwertigen allphatischen Kohlenwasserstoffrest darstellt; R¹ und R², die gleich oder verschieden sein können, ein Wasserstoffatom oder einen substituierten oder unsubstituierten einwertigen Kohlenwasserstoffrest darstellen, mit der Maßgabe, daß, falle R³ ein mehrwertiger allphatischer Kohlenwasserstoffrest ist, entweder R¹ oder R² kein Wasserstoffatom ist; und n eine ganze Zahl von 1 bis 6 darstellt;

- (C) oin aprotisches Lösungsmittel, das Ire) von halogenlerten Kohlenwasserstoffen ist; und
- (D) eine Lewissäure, ausgenommen Dialkylaluminiumchlorid und Dialkylborchlorid,

wobel die Komponente (C) eine Dielektrizitätskonstante von nicht weniger als 2,0 bls nicht mehr als 4,0 bei einer Temperatur von 20°C aufweist.

- Verfahren zur Herstellung eines Isobutenpolymers, das endständige funktionalle Gruppen aufweist bei einer Temperatur nicht unterhalb von-100°C bis nicht oberhalb von 0°C aus den folgenden Komponenten als wesentliche Komponenten:
 - (A) ein kationisch polymerisierbares Monomor, das Isobuten enthält;
 - (B) eine organische Verbindung als Initiator/Kettenübertragungsreagens, dargestellt durch die Formel (I)

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wobei X ein Halogenatom, einen RO-Rest oder RCOO-Rest daretellt (wobei R einen einwertigen organischen Rest darstellt); R³ einen mehrwertigen aromatischen Rest oder einen substituierten oder unsubstituierten mehrwertigen allphatischen Kohlenwasserstoffrest darstellt; R¹ und R², die gleich oder verschieden sein können, ein Wasserstoffatom oder einen substituierten oder unsubstituierten einwertigen Kohlenwasserstoffrest darstellen, mit der Maßgabe, daß, falls R³ ein mehrwertiger allphatischer Kohlenwasserstoffrest ist, entweder R¹ oder R² kein Wasserstoffatom ist; und n eine ganze Zahl von 1 bis 6 darstellt;

- (C) ein aprotisches Lösungsmillel, das halogenlerten Kohlenwasserstoff enthält; und
- (D) eine Lewissäure, ausgenommen Dialkylaluminlumchlorld und Dialkylborchlorid,

wobel die Komponento (C) oino Dielektrizitätskonstante von nicht woniger als 3,5 bis nicht mehr als 5,5 bei einor Temperatur von 20°C aufweist.

- Verfahren zur Herstellung eines durch funktionelle Gruppen abgeschlossenen leobutenpolymers gem
 äß Anspruch 1 oder 2, wobel Komponente (C) einen aromalischen Kohlenwasserstoff enth
 ält.
 - Verfahren zur Herstellung eines durch funktionelle Gruppen abgeschlossenen Isobutenpolymers gemäß einem der Ansprüche 1 bis 3, wobel Komponente (C) eine Mischung aus einem aromatischen Kohlenwasserstoff und einem aliphatischen Kohlenwasserstoff ist.
 - Verfahren zur Herstellung eines durch tunktionello Gruppen abgeschlossenen Isobutenpolymers gemäß einem der Ansprüche 1, 3 und 4, webei Komponente (C) eine Dielektrizitätskonstante von nicht weniger als 2,0 bis nicht mohr als 2,5 bei einer Temperatur von 20°C aufweist.
 - Verfahren zur Herstellung eines durch funktionelle Gruppen abgeschlossenen Isobutenpolymers gemäß einem der Ansprüche 3 bis 5, wobei der aromatische Kohlenwassereloff ausgewählt ist aus Benzol, Toluol, Xylol, und Ethylbenzol.
- Verfahren zur Herstellung eines durch funktionelle Gruppen abgeschlossenen Isobulenpolymere gem
 äß einem
 der Ansprüche 1 bie 6, wobei die Konzentration der Komponenie (A) im System bei 5 bie 40 Gew.-% liegt.
 - Verfahren zur Herstellung eines durch funktionelle Gruppen abgeschlossenen Isobutenpolymere gem
 ß einem
 der Ansprüche 1 bis 7, wobei Komponente (B) ausgewählt ist aus p-Dicumylchlorid, m-Dicumylchlorid, p-Dicumylmethoxid, m-Dicumylmethoxid, 1,3,5-Tricumylchlorid und 1,3,5-Tricumylmethoxid.
 - Verfahren zur Herstellung eines durch funktionelle Gruppen abgeschlossenen Isobutenpolymers gamäß einem der Aneprüche 1 bis 8, wobel Komponente (D) Bortrichlorid oder Titantetrachlorid ist.
- 45 10. Verfahren zur Herstellung eines durch funktionelle Gruppen abgeschlossenen leobutenpolymers gamäß einem der Ansprüche 1 bis 9, wobei weiterhin ein Elektronendonor als Komponente (E) zusätzlich zu den Komponenten (A) bis (D) zugegeben wird.
- 11. Vertahren zur Herstellung eines Isobutenpolymers, das endständige Kohlenstoff-Kohlenstoff-Doppelbindungen trägt, umfassend die Zugabe eines Allykrimethylsikans in einer Menge von 0,5 bis 1,5 Mol pro Mol des Rastes X in Komponente (B) vor der Erzougung eines durch funktionelle Gruppen abgeschlossenen Isobutenpolymers gemäß einem der Ansprüche 1 bis 10 oder vor der Deaktivierung der Komponente (D), die der Erzeugung des Isobutenpolymers folgt.
- 12. Verfahren zur Herstellung eines Isobutenpolymers, das endständige Kohlenstoff-Kohlenstoff-Doppelbindungen trägt, umfassend die Zugabe von 1,9-Decadien in einer Menge von 1,5 bis 10,0 Mol pro Mol des Restes X in Komponente (B) vor der Erzeugung eines durch funktionelle Gruppen abgeschlossenen Isobutenpolymers gemäß einem der Ansprüche 1 bis 10 oder vor der Doaktivierung der Komponente (D), die der Erzeugung des Isobuten-

polymers folgl.

Revendications

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- Procédé pour la préparation d'un polymère d'isobutène présentant des groupes fonctionnels aux torminaisons à une température d'au moins -100°C à au plus 0°C à partir des constituants suivants en tant que constituants essentiels:
 - (A) un monomère cationiquement polymorisable contenant de l'isobutène ;
 - (B) un composé organique en lant qu'initiateur/agent de transfert de chaîne représenté par la formule (1) :

$$R^{1} \xrightarrow{C} X)_{p} \tag{1}$$

dans laquelle X roprésente un atome d'halogène, un groupe RO- ou un groupe RCOO- (dans lequel R représente un groupe organique monovalent); R³ représente un groupe atomatique polyvalent ou un groupe hydrocarboné aliphatique polyvalent substitué ou non substitué; R¹ et R² peuvent être identiques ou différents et représentent chacun un atome d'hydrogène ou un groupe hydrocarboné monovalent substitué ou non substitué, à la condition que lorsque R³ est un groupe hydrocarboné aliphatique polyvalent, un parmi R¹ et R² n'est pas un atome d'hydrogène; et n représente un nombre entier de 1 à 6;

- (C) un solvant aprotique exempt d'hydrocarbure halogéné; ot
- (D) un acide de Lewis à l'exclusion du chlorure de dialkylaluminium et du chlorure de dialkylbore.
- dans lequel ledit constituant (C) présente une constante diélectrique d'au moins 2,0 à au plus 4,0 à une lempérature de 20°C.
 - Procédé pour la préparation d'un polymère d'isobutène présentant des groupes fonctionnels aux terminaisons à une température d'au moins -100°C à au plus 0°C à partir des constituants suivants en tant que constituants essentiels;
 - (A) un monomère cationiquement polyméricable contenant de l'isobutène ;
 - (B) un composé organique en tant qu'initiateur/agent de transfert de chaîné représenté par la formule (1):

$$R^{3} \longrightarrow (C \longrightarrow X)_{n}$$
 (1)

dans lequelle X représente un atome d'halogène, un groupe RO- ou un groupe RCOO- (dans lequel R représente un groupe organique monovalent); R³ représente un groupe atomatique polyvalent ou un groupe hydrocarboné aliphatique polyvalent substitué ou non substitué; R¹ et R² peuvent âtre identiques ou différents et représentent chacun un atome d'hydrogène ou un groupe hydrocarboné monovalent substitué ou non substitué, à la condition que lorsque R³ est un groupe hydrocarboné aliphatique polyvalent, un parmi R¹ et R² n'est pas un atomo d'hydrogène; et n représente un nombre ontier de 1 à 8;

- (C) un solvant aprotique contenant un hydrocarbure halogéné; et
- (D) un acide de Lewis à l'exclusion du chlorure de dialkylaluminium et du chlorure de dialkylbore.

dans loquel ledit constituant (C) présente une constante diélectrique d'au moins 3,5 à au plue 5,5 à une température

de 20°C.

- Procédé pour la préparation d'un polymère d'isobutène terminé par des groupes fonctionnels selon la revendication 1 ou 2, dans lequel ledit constituant (C) contient un hydrocarbure aromatique.
- 4. Procédé pour la préparation d'un polymère d'isobutène terminé par des groupes fonctionnels selon l'une quelconque des revendications 1 à 3, dans lequel ledit constituant (C) est un mélange d'un hydrocarbure aromatique et d'un hydrocarbure aliphatique.
- 70 5. Procédé pour la préparation d'un polymère d'isobutène terminé par des groupes fonctionnels solon l'une quelconque des revendications 1, 3 et 4, dans lequel ladit constituent (C) présente une constante diélectrique d'au moins 2,0 à au plus 2,5 à une température de 20°C.
- 5. Procédé pour la préparation d'un polymère d'isobutène terminé par des groupes fonctionnels selon l'une quelconque des revendications 3 à 5, dans lequel ledit hydrocarbure aromatique est choisi parmi le benzène, le toluène, le xylòno et l'éthylbenzène.
- Procédé pour la préparation d'un polymère d'isobutène terminé par des groupes fonctionnels selon l'une quelconque des revendications 1 à 8, dans lequel la concentration dudit constituant (A) dans le système est do 5 à 40 % en poids.
 - 8. Procédé pour la préparation d'un polymère d'isobutène terminé par des groupes fonctionnels selon l'une quelconque des revendications 1 à 7, dans lequet ledit constituant (B) est choisi parmi le chlorure de p-dicumylo, le chlorure de m-dicumyle, le p-dicumylméthoxyde, le m-dicumylméthoxyde, le chlorure de 1,3,5-tricumyle et le 1,3,5-tricumyle et le 1,3,5-tricumyle.
 - Procédé pour la préparation d'un polymère d'isobutène terminé par des groupes fonctionnels salon l'une quelconque des revendications 1 à 8, dans lequel ledit constituant (D) est le trichlorure de bore ou le tétrachlorure de titane.
- 30 10. Procédé pour la préparation d'un polymère d'isobulène terminé par des groupes (onctionnéle selon l'une quelconque des revendications 1 à 9, dans lequel un donneur d'électrons est en outre ajouté en tant que constituent (E) en plus desdits constituents (A) à (D).
- 11. Procédé pour la préparation d'un polymère d'isobutène présentant des doubles liaisons carbone-carbone aux terminaisons, lequel comprend l'addition d'un allyltriméthylsilane dans une quantité de 0,5 à 1,5 mole par mole du groupe X dans ledit constituant (B) avant la formation d'un polymère d'isobutène terminé par des groupes tonctionnels selon l'une quelconque dos revendications 1 à 10 ou avant la désactivation dudit constituant (D) suivant la formation dudit polymère d'isobutène.
- 40 12. Procédé pour la préparation d'un polymère d'isobutène présentant des doublos liaisons carbone-carbone aux terminalsone, lequel comprend l'addition de 1,9-décadiène dans une quantité de 1,5 à 10,0 mole par mole du groupe X dans lodit constituant (B) avant la formation d'un polymère d'isobutène terminé par des groupes fonctionnels selon l'une quelconque des revendications 1 à 10 ou avant la désactivation dudit constituant (D) suivant la formation dudit polymère d'isobutène.

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